Practical Scientific Computing in Python A Workbook

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Introduction

This document contains a set of small problems, drawn from many different fields, meant to illustrate commonly useful techniques for using Python in scientific computing.

All problems are presented in a similar fashion: the task is explained including any necessary mathematical background and a 'code skeleton' is provided that is meant to serve as a starting point for the solution of the exercise. In some cases, some example output of the expected solution, figures or additional hints may be provided as well.

The accompanying source download for this workbook contains the complete solutions, which are not part of this document for the sake of brevity.

For several examples, the provided skeleton contains pre-written tests which validate the correctness of the expected answers. When you have completed the exercise successfully, you should be able to run it from within IPython and see something like this (illustrated using a trapezoidal rule problem, whose solution is in the file trapezoid.py):

```
In [7]: run trapezoid.py
....
Ran 4 tests in 0.003s
```

This message tells you that 4 automatic tests were successfully executed. The idea of including automatic tests in your code is a common one in modern software development, and Python includes in its standard library two modules for automatic testing, with slightly different functionality: unittest and doctest. These tests were written using the unittest system, whose complete documentation can be found here: http://docs.python.org/lib/module-unittest.html.

Other exercises will illustrate the use of the doctest system, since it provides complementary functionality.

Simple non-numerical problems

1. Sorting quickly with QuickSort

Quicksort is one of the best known, and probably the simplest, fast algorithm for sorting n items. It is fast in the sense that it requires on average $\mathcal{O}(n \log n)$ comparisons instead of $\mathcal{O}(n^2)$, although a naive implementation does have quadratic worst-case behavior.

The algorithm uses a simple divide and conquer strategy, and its implementation is naturally recursive. Its basic steps are:

- (1) Pick an element from the list, called the pivot p (any choice works).
- (2) Select from the rest of the list those elements smaller and those greater than the pivot, and store them in separate lists S and G.
- (3) Recursively apply the algorithm to S and G. The final result can be written as $\sigma(S)$ + $[p] + \sigma(G)$, where σ represents the sorting operation, + indicates list concatenation and [p] is the list containing the pivot as its single element.

The listing 2.1 contains a skeleton with no implementation but with tests already written (in the form of *unit tests*, as described in the introduction).

Listing 2.1

```
"""Simple quicksort implementation."""
def qsort(lst):
    """Return a sorted copy of the input list."""
    raise NotImplementedError
if __name__ == '__main__':
    from unittest import main, TestCase
    import random
    class qsortTestCase(TestCase):
        def test_sorted(self):
            seq = range(10)
            sseq = qsort(seq)
            self.assertEqual(seq, sseq)
        def test_random(self):
            tseq = range(10)
            rseq = range(10)
            random.shuffle(rseq)
            sseq = qsort(rseq)
            self.assertEqual(tseq,sseq)
    main()
```

Hints.

• Python has no particular syntactic requirements for implementing recursion, but it does have a maximum recursion depth. This value can be queried

via the function sys.getrecursionlimit(), and it can be changed with sys.setrecursionlimit(new_value).

- Like in all recursive problems, don't forget to implement an exit condition!
- If L is a list, the call len(L) provides its length.

2. Dictionaries for counting words

A common task in text processing is to produce a count of word frequencies. While NumPy has a builtin histogram function for doing numerical histograms, it won't work out of the box for couting discrete items, since it is a binning histogram for a range of real values.

But the Python language provides very powerful string manipulation capabilities, as well as a very flexible and efficiently implemented builtin data type, the *dictionary*, that makes this task a very simple one.

In this problem, you will need to count the frequencies of all the words contained in a compressed text file supplied as input.

The listing 2.2 contains a skeleton for this problem, with XXX marking various places that are incomplete.

Listing 2.2

```
#!/usr/bin/env python
"""Word frequencies - count word frequencies in a string."""
def word_freq(text):
    """Return a dictionary of word frequencies for the given text."""
    # XXX you need to write this
def print_vk(lst):
    """Print a list of value/key pairs nicely formatted in key/value order."""
    # Find the longest key: remember, the list has value/key paris, so the key
    # is element [1], not [0]
    longest_key = max(map(lambda x: len(x[1]), lst))
    # Make a format string out of it
    fmt = '%'+str(longest_key)+'s -> %s'
    # Do actual printing
    for v,k in 1st:
       print fmt % (k,v)
def freq_summ(freqs, n=10):
    """Print a simple summary of a word frequencies dictionary.
      - freqs: a dictionary of word frequencies.
    Optional inputs:
      - n: the number of items to print"""
    words, counts = # XXX look at the keys and values methods of dicts
    # Sort by count
    items = # XXX think of a list, look at zip() and think of sort()
    print 'Number of words:',len(freqs)
   print
    print '%d least frequent words:' % n
```

```
print_vk(items[:n])
print
print '%d most frequent words:' % n
print_vk(items[-n:])

if __name__ == '__main__':
    text = # XXX
# You need to read the contents of the file HISTORY.gz. Do NOT unzip it
# manually, look at the gzip module from the standard library and the
# read() method of file objects.
freqs = word_freq(text)
freq_summ(freqs, 20)
```

Hints.

- The print_vk function is already provided for you as a simple way to summarize your results.
- You will need to read the compressed file <code>HISTORY.gz</code>. Python has facilities to do this without having to manually uncompress it.
- Consider 'words' simply the result of splitting the input text into a list, using any form of whitespace as a separator. This is obviously a very naïve definition of 'word', but it shall suffice for the purposes of this exercise.
- Python strings have a .split() method that allows for very flexible splitting. You can easily get more details on it in IPython:

The complete set of methods of Python strings can be viewed by hitting the TAB key in IPython after typing 'a.', and each of them can be similarly queried with the '?' operator as above. For more details on Python strings and their companion sequence types, see http://docs.python.org/lib/typesseq.html.

Working with files, the internet, and numpy arrays

This section is a general overview to show how easy it is to load and manipulate data on the file system and over the web using python's built in data structures and numpy arrays. The goal is to exercise basic programming skills like building filename or web addresses to automate certain tasks like loading a series of data files or downloading a bunch of related files off the web, as well as to illustrate basic numpy and pylab skills.

1. Loading and saving ASCII data

The simplest file format is a plain text ASCII file of numbers. Although there are many better formats out there for saveing and loading data, this format is extremely common because it has the advantages of being human readable, and thus will survive the test of time as the *en vogue* programming languages, analysis applications and data formats come and go, it is easy to parse, and it is supported by almost all languages and applications.

In this exercise we will create a data set of two arrays, the first one regularly sampled time t from 0..2 seconds with 20 ms time step, and the second one an array v of sinusoidal voltages corrupted by some noise. Let's assume the sine wave has amplitude 2 V, frequency 10 Hz, and zero mean Gaussian distributed white noise with standard deviation 0.5 V. Your task is to write two scripts.

The first script should create the vectors t and v, plot the time series of t versus v, save them in a two dimensional numpy array X, and then dump the array X to a plain text ASCII file called 'noisy_sine.dat'. The file will look like (not identical because of the noise)

Here is the exercise skeleton of the script to create and plot the data file

Listing 3.1

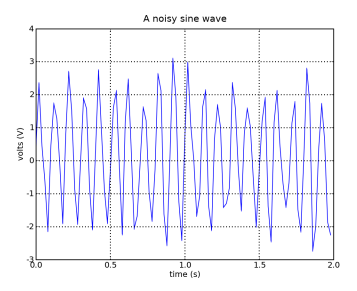


FIGURE 1. A 10 Hz sine wave corrupted by noise

```
# save the output file as ASCII; see the pylab command save
XXX

# plot the arrays t vs v and label the x-axis, y-axis and title save
# the output figure as noisy_sine.png. See the pylab commands plot,
# xlabel, ylabel, grid, show
XXX
```

and the graph will look something like Figure 1

The second part of this exercise is to write a script which loads data from the data file into an array X, extracts the columns into arrays t and v, and computes the RMS (root-mean-square) intensity of the signal using the load command.

2. Working with CSV files

The CSV (Comma Separated Value) file specification is also an ASCII based, human readable format, but it is more powerful than simple flat ASCII files including headers, escape sequences and arbitrary delimiters like TAB, SPACE or COMMA. It is a widely used interchange format for sharing data between operating systems and programs like Excel, Matlab and statistical analysis packages.

A typical CSV file will be a mix of different data types: integers, floating point numbers, dates and strings. Of course, all of these are strings in the file, since all text files are made up of strings, but the data is typically representing some other numeric or date type. Python has very good support for handling different data types, so you don't need to try to force your data to look like a multi dimensional array of floating point numbers if this is not the natural way to describe your data. numpy provides a generalization of the array data structure we used above called record arrays, which allow to store data in a conceptual model similar to a database or spreadsheet: several named fields (eg 'date', 'weight', 'height', 'age') with different types (eg datetime.date, float, float, int).

In the example below, we will download some CSV files from Yahoo Financial web pages and load them into numpy record arrays for analysis and visualization. Go to http://finance.yahoo.com and enter a stock symbol in the entry boc labeled "Get Quotes". I will use 'SPY' which is an index fund that tracks the S&P 500. In the left menu bar, there is

an entry called "Historical Prices" which will take you to a page where you can download the price history of your stock. Near the bottom of this page you should see a "Download To Spreadsheet" link – instead of clicking on it, right click it and choose "Copy Link Location" and paste this into a python script or ipython session as a string named url. Eg, for SPY page better

```
url = 'http://ichart.finance.yahoo.com/table.csv?' +\
   's=SPY&d=9&e=20&f=2007&g=d&a=0&b=29&c=1993&ignore=.csv'
```

I've broken the url into two strings so they will fit on the page. If you spend a little time looking at this pattern, you can probably figure out what is going on. The URL is encoding the information about the stock, the variable s for the stock ticker, d for the latest month, e for the latest day, f for the latest year, c for the start year, and so on (similarly a, b, and c for the start month, day and year). This is handy to know, because below we will write some code to automate some downloads for a stock universe.

One of the great things about python is it's "batteries included" standard library, which includes support for dates, csv files and internet downloads. The example interactive session below shows how in just a few lines of code using python's urllib for retrieving information from the internet, and matplotlib's csv2rec function for loading numpy record arrays, we are ready to get to work analyzing some web based data. Comments have been added to a copy-and-paste from the interactive session

```
# import a couple of libraries we'll be needing
In [23]: import urllib
In [24]: import matplotlib.mlab as mlab
# this is the CSV file we'll be downloading
In [25]: url = 'http://ichart.finance.yahoo.com/table.csv?' +\
   's=SPY&d=9&e=20&f=2007&g=d&a=0&b=29&c=1993&ignore=.csv'
# this will grab that web file and save it as 'SPY.csv' on our local
# filesystem
In [27]: urllib.urlretrieve(url, 'SPY.csv')
Out[27]: ('SPY.csv', <httplib.HTTPMessage instance at 0x2118210>)
# here we use the UNIX command head to peak into the file, which is
# a comma separated and contains various types, dates, ints, floats
In [28]: !head SPY.csv
Date, Open, High, Low, Close, Volume, Adj Close
2007-10-19,153.09,156.48,149.66,149.67,295362200,149.67
2007-10-18, 153.45, 154.19, 153.08, 153.69, 148367500, 153.69
2007-10-17, 154.98, 155.09, 152.47, 154.25, 216687300, 154.25
2007-10-16, 154.41, 154.52, 153.47, 153.78, 166525700, 153.78
2007-10-15, 156.27, 156.36, 153.94, 155.01, 161151900, 155.01
2007-10-12,155.46,156.35,155.27,156.33,124546700,156.33
2007-10-11,156.93,157.52,154.54,155.47,233529100,155.47
2007-10-10, 156.04, 156.44, 155.41, 156.22, 101711100, 156.22
2007-10-09, 155.60, 156.50, 155.03, 156.48, 94054300, 156.48
# csv2rec will import the file into a numpy record array, inspecting
# the columns to determine the correct data type
In [29]: r = mlab.csv2rec('SPY.csv')
# the dtype attribute shows you the field names and data types.
# 04 is a 4 byte python object (datetime.date), f8 is an 8 byte
# float, i4 is a 4 byte integer and so on. The > and < symbols
# indicate the byte order of multi-byte data types, eq biq endian or
```

```
# little endian, which is important for cross platform binary data
# storage
In [30]: r.dtype
Out[30]: dtype([('date', '|04'), ('open', '>f8'), ('high', '>f8'),
('low', '>f8'), ('close', '>f8'), ('volume', '>i4'), ('adj_close',
'>f8')])
# Each of the columns is stored as a numpy array, but the types are
# preserved. Eg, the adjusted closing price column adj_close is a
# floating point type, and the date column is a python datetime.date
In [31]: print r.adj_close
[ 149.67 153.69 154.25 ...,
                                34.68
                                        34.61
                                                34.361
In [32]: print r.date
[2007-10-19 00:00:00 2007-10-18 00:00:00 2007-10-17 00:00:00 ...,
1993-02-02 00:00:00 1993-02-01 00:00:00 1993-01-29 00:00:00]
```

For your exercise, you'll elaborate on the code here to do a batch download of a number of stock tickers in a defined stock universe. Define a function fetch_stock (ticker) which takes a stock ticker symbol as an argument and returns a numpy record array. Select the rows of the record array where the date is greater than 2003-01-01 and plot the returns $(p-p_0)/p_0$ where p are the prices and p_0 is the initial price. by date for each stock on the same plot. Create a legend for the plot using the matplotlib legend command, and print out a sorted list of final returns (eg assuming you bought in 2003 and held to the present) for each stock. Here is the exercise skeleton:

Listing 3.2

```
0.00
Download historical pricing record arrays for a universe of stocks
from Yahoo Finance using urllib. Load them into numpy record arrays
using matplotlib.mlab.csv2rec, and do some batch processing -- make
date vs price charts for each one, and compute the return since 2003
for each stock. Sort the returns and print out the tickers of the 4
biggest winners
0.00
import os, datetime, urllib
import matplotlib.mlab as mlab # contains csv2rec
import numpy as npy
import pylab as p
def fetch_stock(ticker):
    download the CSV file for stock with ticker and return a numpy
    record array. Save the CSV file as TICKER.csv where TICKER is the
    stock's ticker symbol.
    Extra credit for supporting a start date and end date, and
    checking to see if the file already exists on the local file
    system before re-downloading it
    11 11 11
    fname = '%s.csv'%ticker
    url = XXX # create the url for this ticker
    # the os.path module contains function for checking whether a file
    # exists, and fetch it if not
    XXX
```

```
# load the CSV file intoo a numpy record array
    r = XXX
    # note that the CSV file is sorted most recent date first, so you
    # will probably want to sort the record array so most recent date
    # is last
    XXX
    return r
tickers = 'INTC', 'MSFT', 'YHOO', 'GOOG', 'GE', 'WMT', 'AAPL'
# we want to compute returns since 2003, so define the start date as a
# datetime.datetime instance
startdate = XXX
# we'll store a list of each return and ticker for analysis later
data = [] # a list of (return, ticker) for each stock
fig = p.figure()
for ticker in tickers:
   print 'fetching', ticker
    r = fetch_stock(ticker)
    # select the numpy records where r.date>=startdatre use numpy mask
    # indexing to restrict r to just the dates > startdate
   r = XXX
    price = XXX  # set price equal to the adjusted close
    returns = XXX \# return is the (price-p0)/p0
                  # store the data
    XXX
    # plot the returns by date for each stock using pylab.plot, adding
    # a label for the legend
   XXX
# use pylab legend command to build a legend
XXX
# now sort the data by returns and print the results for each stock
XXX
# show the figures
p.show()
```

The graph will look something like Figure 2.

3. Loading and saving binary data

ASCII is bloated and slow for working with large arrays, and so binary data should be used if performance is a consideration. To save an array X in binary form, you can use the numpy tostring method

```
In [16]: import numpy
# create some random numbers
In [17]: x = numpy.random.rand(5,2)
```

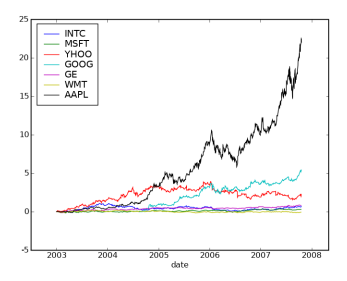


Figure 2. Returns for a universe of stocks since 2003

```
In [19]: print x
[[ 0.56331918  0.519582 ]
 [ 0.22685429  0.18371135]
 [ 0.19384767  0.27367054]
 [ 0.35935445  0.95795884]
 [ 0.37646642  0.14431089]]
# save it to a data file in binary
In [20]: x.tofile(file('myx.dat', 'wb'))
# load it into a new array
In [21]: y = numpy.fromfile(file('myx.dat', 'rb'))
# the shape is not preserved, so we will have to reshape it
In [22]: print y
[ 0.56331918  0.519582
                          0.22685429 0.18371135 0.19384767
0.27367054
  0.35935445 0.95795884 0.37646642 0.14431089]
In [23]: y.shape
Out[23]: (10,)
# restore the original shape
In [24]: y.shape = 5, 2
In [25]: print y
[[ 0.56331918  0.519582 ]
 [ 0.22685429  0.18371135]
 [ 0.19384767  0.27367054]
 [ 0.35935445  0.95795884]
 [ 0.37646642  0.14431089]]
```

The advantage of numpy tofile and fromfile over ASCII data is that the data storage is compact and the read and write are very fast. It is a bit of a pain that that meta ata like array datatype and shape are not stored. In this format, just the raw binary numeric data is stored, so you will have to keep track of the data type and shape by other means. This is a good solution if you need to port binary data files between different packages, but if you know you will always be working in python, you can use the python pickle function to preserve all metadata (pickle also works with all standard python data types, but has the disadvantage that other programs and applications cannot easily read it)

```
# create a 6,3 array of random integers
In [36]: x = (256*numpy.random.rand(6,3)).astype(numpy.int)
In [37]: print x
[[173 38
           2]
 [243 207 155]
 [127 62 140]
 [ 46 29 98]
 [ 0 46 156]
 [ 20 177 36]]
# use pickle to save the data to a file myint.dat
In [38]: import cPickle
In [39]: cPickle.dump(x, file('myint.dat', 'wb'))
# load the data into a new array
In [40]: y = cPickle.load(file('myint.dat', 'rb'))
# the array type and share are preserved
In [41]: print y
[[173 38
            2]
[243 207 155]
 [127 62 140]
 [ 46 29 98]
 [ 0 46 156]
 [ 20 177 36]]
```

Elementary Numerics

1. Wallis' slow road to π

Wallis' formula is an infinite product that converges (slowly) to π :

(1)
$$\pi = \prod_{i=1}^{\infty} \frac{4i^2}{4i^2 - 1}.$$

The listing 4.1 contains a skeleton with no implementation but with some plotting commands already inserted, so that you can visualize the convergence rate of this formula as more terms are kept.

Listing 4.1

```
#!/usr/bin/env python
"""Simple demonstration of Python's arbitrary-precision integers."""
# We need exact division between integers as the default, without manual
# conversion to float b/c we'll be dividing numbers too big to be represented
# in floating point.
from __future__ import division
from decimal import Decimal
def pi(n):
    """Compute pi using n terms of Wallis' product.
    Wallis' formula approximates pi as
    pi(n) = 2 \prod_{i=1}^{n} \frac{4i^2}{4i^2-1}."""
    XXX
# This part only executes when the code is run as a script, not when it is
# imported as a library
if __name__ == '__main__':
    # Simple convergence demo.
    # A few modules we need
    import pylab as P
    import numpy as N
    # Create a list of points 'nrange' where we'll compute Wallis' formula
    nrange = XXX
    # Make an array of such values
    XXX = iqw
    # Compute the difference against the value of pi in numpy (standard
```

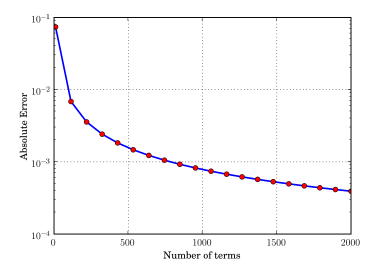


FIGURE 1. Convergence rate for Wallis' infinite product approximation to π .

```
# 16-digit value)
diff = XXX

# Make a new figure and build a semilog plot of the difference so we can
# see the quality of the convergence

P.figure()
# Line plot with red circles at the data points
P.semilogy(nrange,diff,'-o',mfc='red')

# A bit of labeling and a grid
P.title(r"Convergence of Wallis' product formula for pi")
P.xlabel('Number of terms')
P.ylabel(r'|Error}|')
P.grid()

# Display the actual plot
P.show()
```

After running the script successfully, you should obtain a plot similar to Figure 1.

2. Trapezoidal rule

In this exercise, you are tasked with implementing the simple trapezoid rule formula for numerical integration. If we want to compute the definite integral

(2)
$$\int_{a}^{b} f(x)dx$$

we can partition the integration interval [a, b] into smaller subintervals, and approximate the area under the curve for each subinterval by the area of the trapezoid created by linearly interpolating between the two function values at each end of the subinterval. This is graphically illustrated in Figure 2, where the blue line represents the function f(x) and the red line represents the successive linear segments.

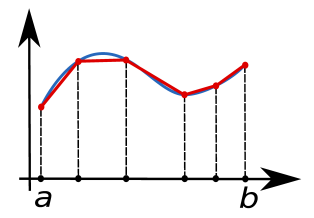


FIGURE 2. Illustration of the composite trapezoidal rule with a non-uniform grid (Image credit: Wikipedia).

The area under f(x) (the value of the definite integral) can thus be approximated as the sum of the areas of all these trapezoids. If we denote by x_i (i = 0, ..., n, with $x_0 = a$ and $x_n = b$) the abscissas where the function is sampled, then

(3)
$$\int_{a}^{b} f(x)dx \approx \frac{1}{2} \sum_{i=1}^{n} (x_{i} - x_{i-1}) (f(x_{i}) + f(x_{i+1})).$$

The common case of using equally spaced abscissas with spacing h = (b - a)/n reads simply

(4)
$$\int_{a}^{b} f(x)dx \approx \frac{h}{2} \sum_{i=1}^{n} (f(x_i) + f(x_{i+1})).$$

One frequently receives the function values already precomputed, $y_i = f(x_i)$, so equation (3) becomes

(5)
$$\int_{a}^{b} f(x)dx \approx \frac{1}{2} \sum_{i=1}^{n} (x_{i} - x_{i-1}) (y_{i} + y_{i-1}).$$

Listing 4.2 contains a skeleton for this problem, written in the form of two incomplete functions and a set of automatic tests (in the form of *unit tests*, as described in the introduction).

Listing 4.2

```
#!/usr/bin/env python
"""Simple trapezoid-rule integrator."""

import numpy as N

def trapz(x, y):
    """Simple trapezoid integrator for sequence-based innput.

Inputs:
    - x,y: arrays of the same length.

Output:
    - The result of applying the trapezoid rule to the input, assuming that y[i] = f(x[i]) for some function f to be integrated.

Minimally modified from matplotlib.mlab."""

raise NotImplementedError
```

```
def trapzf(f,a,b,npts=100):
    """Simple trapezoid-based integrator.
    Inputs:
     - f: function to be integrated.
      - a,b: limits of integration.
    Optional inputs:
      - npts(100): the number of equally spaced points to sample f at, between
      a and b.
    Output:
      - The value of the trapezoid-rule approximation to the integral."""
    # you will need to apply the function f to easch element of the
    # vector x. What are several ways to do this? Can you profile
    # them to see what differences in timings result for long vectors
    # x?
    raise NotImplementedError
if __name__ == '__main__':
    # Simple tests for trapezoid integrator, when this module is called as a
    # script from the command line.
    import unittest
    import numpy.testing as ntest
    def square(x): return x**2
    class trapzTestCase(unittest.TestCase):
        def test_err(self):
            self.assertRaises(ValueError, trapz, range(2), range(3))
        def test_call(self):
            x = N.linspace(0,1,100)
            y = N.array(map(square, x))
            ntest.assert_almost_equal(trapz(x, y),1./3,4)
    class trapzfTestCase(unittest.TestCase):
        def test_square(self):
            ntest.assert_almost_equal(trapzf(square, 0, 1), 1./3, 4)
        def test_square2(self):
            ntest.assert_almost_equal(trapzf(square,0,3,350),9.0,4)
    unittest.main()
```

In this exercise, you'll need to write two functions, trapz and trapzf. trapz applies the trapezoid formula to pre-computed values, implementing equation (5), while trapzf takes a function f as input, as well as the total number of samples to evaluate, and computes eq. (4).

3. Newton's method

Consider the problem of solving for t in

(6)
$$\int_{0}^{t} f(s)ds = u$$

where f(s) is a monotonically increasing function of s and u > 0.

This problem can be simply solved if seen as a root finding question. Let

(7)
$$g(t) = \int_0^t f(s)ds - u,$$

then we just need to find the root for g(t), which is guaranteed to be unique given the conditions above.

The SciPy library includes an optimization package that contains a Newton-Raphson solver called scipy.optimize.newton. This solver can optionally take a known derivative for the function whose roots are being sought, and in this case the derivative is simply

(8)
$$\frac{dg(t)}{dt} = f(t).$$

For this exercise, implement the solution for the test function

$$f(t) = t\sin^2(t),$$

using

$$u = \frac{1}{4}.$$

The listing 4.3 contains a skeleton that includes for comparison the correct numerical value.

Listing 4.3

```
#!/usr/bin/env python
"""Root finding using SciPy's Newton's method routines.
from math import sin
import scipy, scipy.integrate, scipy.optimize
quad = scipy.integrate.quad
newton = scipy.optimize.newton
# test input function f(t): t * sin^2(t)
def f(t): XXX
# Use u=0.25
def g(t): XXX
# main
tguess = 10.0
print "Solution using the numerical integration technique"
t0 = newton(g,tguess,f)
print "t0, g(t0) = ", t0, g(t0)
print
print "To six digits, the answer in this case is t==1.06601."
```

Linear algebra

Like matlab, numpy and scipy have support for fast linear algebra built upon the highly optimized LAPACK, BLAS and ATLAS fortran linear algebra libraries. Unlike Matlab, in which everything is a matrix or vector, and the '*' operator always means matrix multiple, the default object in numpy is an array, and the '*' operator on arrays means element-wise multiplication.

Instead, numpy provides a matrix class if you want to do standard matrix-matrix multiplication with the '*' operator, or the dot function if you want to do matrix multiplies with plain arrays. The basic linear algebra functionality is found in numpy.linalg

```
In [1]: import numpy as npy
In [2]: import numpy.linalg as linalg
# X and Y are arrays
In [3]: X = npy.random.rand(3,3)
In [4]: Y = npy.random.rand(3,3)
# * operator is element wise multiplication, not matrix matrix
In [5]: print X*Y
[[ 0.00973215  0.18086148  0.05539387]
[ 0.00817516  0.63354021  0.2017993 ]
# the dot function will use optimized LAPACK to do matrix-matix
# multiply
In [6]: print npy.dot(X, Y)
# the matrix class will create matrix objects that support matrix
\# multiplication with *
In [7]: Xm = npy.matrix(X)
In [8]: Ym = npy.matrix(Y)
In [9]: print Xm*Ym
# the linalg module provides functions to compute eigenvalues,
# determinants, etc. See help(linalg) for more info
In [10]: print linalg.eigvals(X)
                  [ 1.46131600+0.j
```

1. Glass Moiré Patterns

When a random dot pattern is scaled, rotated, and superimposed over the original dots, interesting visual patterns known as Glass Patterns emerge¹ In this exercise, we generate random dot fields using numpy's uniform distribution function, and apply transformations to the random dot field using a scale S and rotation R matrix $X_2 = SRX_1$.

If the scale and rotation factors are small, the transformation is analogous to a single step in the numerical solution of a 2D ODE, and the plot of both X_1 and X_2 will reveal the structure of the vector field flow around the fixed point (the invariant under the transformation); see for example the *stable focus*, aka *spiral*, in Figure 1.

The eigenvalues of the tranformation matrix $\mathbf{M} = \mathbf{SR}$ determine the type of fix point: center, stable focus, saddle node, etc.... For example, if the two eigenvalues are real but differing in signs, the fixed point is a saddle node. If the real parts of both eigenvalues are negative and the eigenvalues are complex, the fixed point is a stable focus. The complex part of the eigenvalue determines whether there is any rotation in the matrix transformation, so another way to look at this is to break out the scaling and rotation components of the transformation \mathbf{M} . If there is a rotation component, then the fixed point will be a center or a focus. If the scaling components are both one, the rotation will be a center, if they are both less than one (contraction), it will be a stable focus. Likewise, if there is no rotation component, the fixed point will be a node, and the scaling components will determine the type of node. If both are less than one, we have a stable node, if one is greater than one and the other less than one, we have a saddle node.

Listing 5.1

```
0.00
Moire patterns from random dot fields
http://en.wikipedia.org/wiki/Moir%C3%A9_pattern
See L. Glass. 'Moire effect from random dots' Nature 223, 578580 (1969).
from numpy import cos, sin, pi, matrix
import numpy as npy
import numpy.linalg as linalg
from pylab import figure, show
def csqrt(x):
    'sqrt func that handles returns sqrt(x) \neq for x<0'
    XXX
def myeig(M):
    compute eigen values and eigenvectors analytically
    Solve quadratic:
      lamba^2 - tau*lambda + Delta = 0
    where tau = trace(M) and Delta = Determinant(M)
    Return value is lambda1, lambda2
    ....
    XXX
\# 2000 random x,y points in the interval[-0.5 ... 0.5]
X1 = XXX
```

¹L. Glass. 'Moiré effect from random dots' Nature 223, 578580 (1969).

```
name = 'saddle'
\#sx, sy, angle = XXX
#name = 'center'
\#sx, sy, angle = XXX
name = 'spiral' #stable focus
sx, sy, angle = XXX
theta = angle \star pi/180. # the rotation in radians
# the scaling matrix
# | sx 0 |
# | 0 sy |
S = XXX
# the rotation matrix
# | cos(theta) -sin(theta) |
# | sin(theta) cos(theta) |
R = XXX
# the transformation is the matrix product of the scaling and rotation
M = XXX
# compute the eigenvalues using numpy linear algebra
print 'numpy eigenvalues', XXX
# compare with the analytic values from myeig
print 'analytic eigenvalues', myeig(M)
\mbox{\tt\#} transform X1 by the matrix \mbox{\tt M}
X2 = XXX
# plot the original X1 as green dots and the transformed X2 as red
# dots
XXX
```

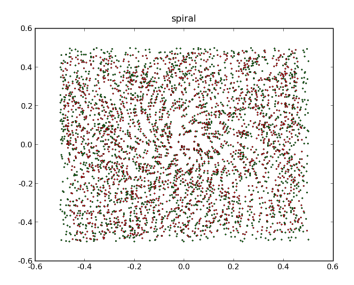


FIGURE 1. Glass pattern showing a stable focus

Signal processing

numpy and scipy provide many of the essential tools for digital signal processing. scipy.signal provides basic tools for digital filter design and filtering (eg Butterworth filters), a linear systems toolkit, standard waveforms such as square waves, and saw tooth functions, and some basic wavelet functionality. scipy.fftpack provides a suite of tools for Fourier domain analysis, including 1D, 2D, and ND discrete fourier transform and inverse functions, in addition to other tools such as analytic signal representations via the Hilbert transformation (numpy.fft also provides basic FFT functions). pylab provides Matlab compatible functions for computing and plotting standard time series analyses, such as historgrams (hist), auto and cross correlations (acorr and xcorr), power spectra and coherence spectra (psd, csd, cohere and specgram).

1. Convolution

The output of a linear system is given by the convolution of its impulse response function with the input. Mathematically

(9)
$$y(t) = \int_0^t x(\tau)r(t-\tau)d\tau$$

This fundamental relationship lies at the heart of linear systems analysis. It is used to model the dynamics of calcium buffers in neuronal synapses, where incoming action potentials are represented as Dirac δ -functions and the calcium stores are represented with a response function with multiple exponential time constants. It is used in microscopy, in which the image distortions introduced by the lenses are *deconvolved* out using a measured point spread function to provide a better picture of the true image input. It is essential in structural engineering to determine how materials respond to shocks.

The impulse response function r is the system response to a pulsatile input. For example, in Figure 1 below, the response function is the sum of two exponentials with different time constants and signs. This is a typical function used to model synaptic current following a neuronal action potential. The figure shows three δ inputs at different times and with different amplitudes. The corresponsing impulse response for each input is shown following it, and is color coded with the impulse input color. If the system response is linear, by definition, the response to a sum of inputs is the sum of the responses to the individual inputs, and the lower panel shows the sum of the responses, or equivalently, the convolution of the impulse response function with the input function.

In Figure 1, the summing of the impulse response function over the three inputs is conceptually and visually easy to understand. Some find the concept of a convolution of an impulse response function with a continuous time function, such as a sinusoid or a noise process, conceptually more difficult. It shouldn't be. By the *sampling theorem*, we can represent any finite bandwidth continuous time signal as the sum of Dirac- δ functions where the height of the δ function at each time point is simply the amplitude of the signal at that time point. The only requirement is that the sampling frequency be at least as high as the Nyquist frequency, defined as the highest spectral frequency in the signal divided by 2. See Figure 2 for a representation of a delta function sampling of a damped, oscillatory, exponential function.

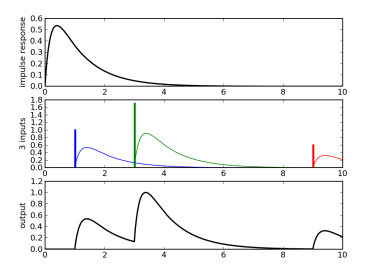


FIGURE 1. The output of a linear system to a series of impulse inputs is equal to the sum of the scaled and time shifted impulse response functions.

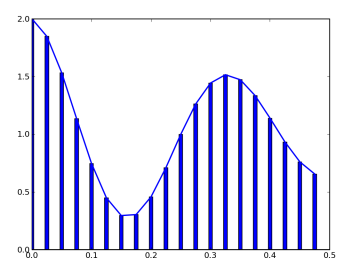


FIGURE 2. Representing a continuous time signal sampled as a sum of delta functions.

In the exercise below, we will convolve a sample from the normal distribution (white noise) with a double exponential impulse response function. Such a function acts as a low pass filter, so the resultant output will look considerably smoother than the input. You can use numpy.convolve to perform the convolution numerically.

We also explore the important relationship that a convolution in the tempoeral (or spatial) domain becomes a multiplication in the spectral domain, which is mathematically much easier to work with.

$$Y = R * X$$

where Y, X, and R are the Fourier transforms of the respective variable in the temporal convolution equation above. The Fourier transform of the impulse response function serves as an

amplitude weighting and phase shifting operator for each frequency component. Thus, we can get deeper insight into the effects of impulse response function r by studying the amplitude and phase spectrum of its transform R. In the example below, however, we simply use the multiplication property to perform the same convolution in Fourier space to confirm the numerical result from numpy.convolve.

```
Listing 6.1
. . .
In signal processing, the output of a linear system to an arbitrary
input is given by the convolution of the impule response function (the
system response to a Dirac-delta impulse) and the input signal.
Mathematically:
 y(t) = \int_0^t x(\tau) r(t-\tau) d\tau
where x(t) is the input signal at time t, y(t) is the output, and r(t)
is the impulse response function.
In this exercise, we will compute investigate the convolution of a
white noise process with a double exponential impulse response
function, and compute the results
  * using numpy.convolve
  * in Fourier space using the property that a convolution in the
    temporal domain is a multiplication in the fourier domain
import numpy as npy
import matplotlib.mlab as mlab
from pylab import figure, show
# build the time, input, output and response arrays
dt = 0.01
t = XXX
               # the time vector from 0..20
Nt = len(t)
def impulse_response(t):
    'double exponential response function'
    return XXX
x = XXX
          # gaussian white noise
# evaluate the impulse response function, and numerically convolve it
# with the input x
r = XXX \# evaluate the impulse function
y = XXX \# convolution of x with r
y = XXX # extract just the length Nt part
```

compute y by applying $F^-1[F(x) * F(r)]$. The fft assumes the signal # is periodic, so to avoid edge artificats, pad the fft with zeros up

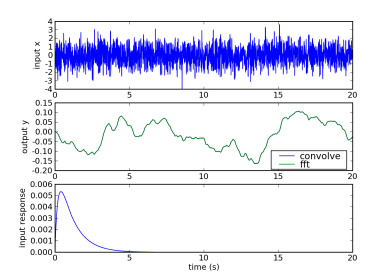


FIGURE 3. Convolution of a white noise process with a double exponential function computed with numpy.fft and numpy.convolve

```
# to the length of r + x do avoid circular convolution artifacts
R = XXX  # the zero padded FFT of r
X = XXX  # the zero padded FFT of x
Y = XXX  # the product of R and S

# now inverse fft and extract the real part, just the part up to
# len(x)
yi = XXX
# plot t vs x, t vs y and yi, and t vs r in three subplots
XXX
show()
```

2. FFT Image Denoising

Convolution of an input with with a linear filter in the termporal or spatial domain is equivalent to multiplication by the fourier transforms of the input and the filter in the spectral domain. This provides a conceptually simple way to think about filtering: transform your signal into the frequency domain, dampen the frequencies you are not interested in by multiplying the frequency spectrum by the desired weights, and then inverse transform the multiplies spectrum back into the original domain. In the example below, we will simply set the weights of the frequencies we are uninterested in (the high frequency noise) to zero rather than dampening them with a smoothly varying function. Although this is not usually the best thing to do, since sharp edges in one domain usually introduce artifacts in another (eg high frequency "ringing"), it is easy to do and sometimes provides satisfactory results.

The image in the upper left panel of Figure 4 is a grayscale photo of the moon landing. There is a banded pattern of high frequency noise polluting the image. In the upper right panel we see the 2D spatial frequency spectrum. The FFT output in scipy is packed with the lower frequencies starting in the upper left, and proceeding to higher frequencies as one moves to the center of the spectrum (this is the most efficient way numerically to fill the output of the FFT algorithm). Because the input signal is real, the output spectrum is complex and symmetrical:

the transformation values beyond the midpoint of the frequency spectrum (the Nyquist frequency) correspond to the values for negative frequencies and are simply the mirror image of the positive frequencies below the Nyquist (this is true for the 1D, 2D and ND FFTs in numpy).

In this exercise we will compute the 2D spatial frequency spectra of the luminance image, zero out the high frequency components, and inverse transform back into the time domain. We can plot the input and output images with the pylab.imshow function, but the images must first be scaled to be withing the 0..1 luminance range. For best results, it helps to amplify the image by some scale factor, and then clip it to set all values greater than one to one. This serves to enhance contrast among the darker elements of the image, so it is not completely dominated by the brighter segments

Listing 6.2

```
#!/usr/bin/env python
"""Image denoising example using 2-dimensional FFT."""
import numpy as N
import pylab as P
import scipy as S
def mag_phase(F):
    """Return magnitude and phase components of spectrum F."""
    # XXX Look at the absolute and angle functions in numpy...
def plot_spectrum(F, amplify=1000):
    """Normalise, amplify and plot an amplitude spectrum."""
    M = # XXX use mag_phase to get the magnitude...
    # XXX Now, rescale M by amplify/maximum_of_M. Numpy arrays can be scaled
    # in-place with ARR *= number. For the max of an array, look for its max
    # method.
    # XXX Next, clip all values larger than one to one. You can set all
    # elements of an array which satisfy a given condition with array indexing
    # syntax: ARR[ARR<VALUE] = NEWVALUE, for example.</pre>
    # Display: this one already works, if you did everything right with M
    P.imshow(M, P.cm.Blues)
# 'main' script
im = # XXX make an image array from the file 'moonlanding.png', using the
     # pylab imread() function. You will need to just extract the red
     # channel from the MxNx4 RGBA matrix to represent the grayscale
     # intensities
F = \# Compute the 2d FFT of the input image. Look for a 2-d FFT in N.dft
# Define the fraction of coefficients (in each direction) we keep
keep_fraction = 0.1
```

```
# XXX Call ff a copy of the original transform. Numpy arrays have a copy
    ...method
# for this purpose.
# XXX Set r and c to be the number of rows and columns of the array. Look for
# the shape attribute...
\# Set to zero all rows with indices between r*keep\_fraction and
# r*(1-keep_fraction):
# Similarly with the columns:
# Reconstruct the denoised image from the filtered spectrum. There's an
# inverse 2d fft in the dft module as well. Call the result im_new
# Show the results.
# The code below already works, if you did everything above right.
P.figure()
P.subplot(221)
P.title('Original image')
P.imshow(im, P.cm.gray)
P.subplot(222)
P.title('Fourier transform')
plot_spectrum(F)
P.subplot (224)
P.title('Filtered Spectrum')
plot_spectrum(ff)
P.subplot (223)
P.title('Reconstructed Image')
P.imshow(im_new, P.cm.gray)
P.show()
```

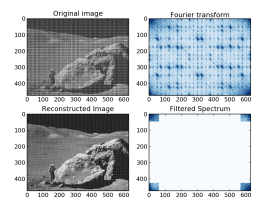


FIGURE 4. High frequency noise filtering of a 2D image in the Fourier domain. The upper panels show the original image (left) and spectral power (right) and the lower panels show the same data with the high frequency power set to zero. Although the input and output images are grayscale, you can provide colormaps to pylab.imshow to plot them in psudo-color

Statistics

R, a statistical package based on S, is viewd by some as the best statistical software on the planet, and in the open source world it is the clear choice for sophisticated statistical analysis. Like python, R is an interpreted language written in C with an interactive shell. Unlike python, which is a general purpose programming language, R is a specialized statistical language. Since python is a excellent glue language, with facilities for providing a transparent interface to FORTRAN, C, C++ and other languages, it should come as no surprise that you can harness R's immense statistical power from python, through the rpy third part extension library.

However, R is not without its warts. As a language, it lacks python's elegance and advanced programming constructs and idioms. It is also GPL, which means you cannot distribute code based upon it unhindered: the code you distribute must be GPL as well (python, and the core scientific extension libraries, carry a more permissive license which support distribution in closed source, proprietary application).

Fortunately, the core tools scientific libraries for python (primarily numpy and scipy.stats) provide a wide array of statistical tools, from basic descriptive statistics (mean, variance, skew, kurtosis, correlation, ...) to hypothesis testing (t-tests, χ -Square, analysis of variance, general linear models, ...) to analytical and numerical tools for working with almost every discrete and continuous statistical distribution you can think of (normal, gamma, poisson, weibull, lognormal, levy stable, ...).

1. Descriptive statistics

The first step in any statistical analysis should be to describe, characterize and importantly, visualize your data. The normal distribution (aka Gaussian or bell curve) lies at the heart of much of formal statistical analysis, and normal distributions have the tidy property that they are completely characterized by their mean and variance. As you may have observed in your interactions with family and friends, most of the world is not normal, and many statistical analyses are flawed by summarizing data with just the mean and standard deviation (square root of variance) and associated signficance tests (eg the T-Test) as if it were normally distributed data.

In the exercise below, we write a class to provide descriptive statistics of a data set passed into the constructor, with class methods to pretty print the results and to create a battery of standard plots which may show structure missing in a casual analysis. Many new programmers, or even experienced programmers used to a proceedural environment, are uncomfortable with the idea of classes, having hear their geekier programmer friends talk about them but not really sure what to do with them. There are many interesting things one can do with classes (aka object oriented programming) but at their hear they are a way of bundling data with methods that operate on that data. The self variable is special in python and is how the class refers to its own data and methods. Here is a toy example

```
In [115]: class MyData:
    ....:    def __init__(self, x):
        self.x = x
    ....:    def sumsquare(self):
        return (self.x**2).sum()
    ....:
    ....:
```

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```
In [116]: nse = npy.random.rand(100)
In [117]: mydata.sumsquare()
Out[117]: 29.6851135284
                                  Listing 7.1
import scipy.stats as stats
from matplotlib.mlab import detrend_linear, load
import numpy
import pylab
XXX = None
class Descriptives:
    a helper class for basic descriptive statistics and time series plots
    def __init__(self, samples):
        self.samples = numpy.asarray(samples)
       self.N = XXX  # the number of samples
       self.median = XXX  # sample median
       self.min = XXX  # sample min
       self.max = XXX
                          # sample max
       self.mean = XXX
                          # sample mean
       self.std = XXX
                          # sample standard deviation
       self.var = XXX
                           # sample variance
       self.skew = XXX  # the sample skewness
       self.kurtosis = XXX # the sample kurtosis
        self.range = XXX  # the sample range max-min
    def __repr__(self):
        Create a string representation of self; pretty print all the
        attributes:
        N, median, min, max, mean, std, var, skew, kurtosis, range,
        descriptives = (
           'N = %d'
                                 % self.N,
           XXX # the rest here
           )
        return '\n'.join(descriptives)
    def plots(self, figfunc, maxlags=20, Fs=1, detrend=detrend_linear, fmt='bo
        ...'):
        ....
       plots the time series, histogram, autocorrelation and spectrogram
        figfunc is a figure generating function, eg pylab.figure
        return an object which stores plot axes and their return
```

```
values from the plots. Attributes of the return object are
        'plot', 'hist', 'acorr', 'psd', 'specgram' and these are the
        return values from the corresponding plots. Additionally, the
        axes instances are attached as c.ax1...c.ax5 and the figure is
        c.fig
        keyword args:
         Fs
                : the sampling frequency of the data
         maxlags : max number of lags for the autocorr
         detrend : a function used to detrend the data for the
         correlation and spectral functions
                : the plot format string
        data = self.samples
        # Here we use a rather strange idiom: we create an empty do
        # nothing class C and simply attach attributes to it for
        # return value (which we carefully describe in the docstring).
        # The alternative is either to return a tuple a,b,c,d or a
        # dictionary {'a':someval, 'b':someotherval} but both of these
        # methods have problems. If you return a tuple, and later
        # want to return something new, you have to change all the
        # code that calls this function. Dictionaries work fine, but
        # I find the client code harder to use d['a'] vesus d.a. The
        # final alternative, which is most suitable for production
        # code, is to define a custom class to store (and pretty
        # print) your return object
        class C: pass
        C = C()
       N = 5
        fig = c.fig = figfunc()
        ax = c.ax1 = fig.add\_subplot(N, 1, 1)
        c.plot = ax.plot(data, fmt)
        # XXX the rest of the plot funtions here
        return c
if __name__=='__main__':
    # load the data in filename fname into the list data, which is a
    # list of floating point values, one value per line. Note you
    # will have to do some extra parsing
    #fname = 'data/nm560.dat' # tree rings in New Mexico 837-1987
   fname = 'data/hsales.dat' # home sales
   for line in file(fname):
        line = line.strip()
```

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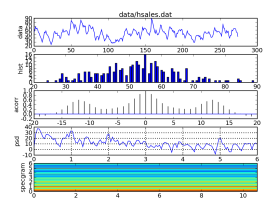


Figure 1.

```
if not line: continue
    # XXX convert to float and add to data here

desc = Descriptives(data)
print desc
c = desc.plots(pylab.figure, Fs=12, fmt='-o')
c.ax1.set_title(fname)
pylab.show()
```

2. Statistical distributions

We explore a handful of the statistical distributions in scipy.stats module and the connections between them. The organization of the distribution functions in scipy.stats is quite elegant, with each distribution providing random variates (rvs), analytical moments (mean, variance, skew, kurtosis), analytic density (pdf, cdf) and survival functions (sf, isf) (where available) and tools for fitting empirical distributions to the analytic distributions (fit).

in the exercise below, we will simulate a radioactive particle emitter, and look at the empirical distribution of waiting times compared with the expected analytical distributions. Our radioative particle emitter has an equal likelihood of emitting a particle in any equal time interval, and emits particles at a rate of 20 Hz. We will discretely sample time at a high frequency, and record a 1 of a particle is emitted and a 0 otherwise, and then look at the distribution of waiting times between emissions. The probability of a particle emission in one of our sample intervals (assumed to be very small compared to the average interval between emissions) is proportional to the rate and the sample interval Δt , ie $p(\Delta t) = \alpha \Delta t$ where α is the emission rate in particles per second.

```
# a uniform distribution [0..1]
In [62]: uniform = scipy.stats.uniform()
# our sample interval in seconds
In [63]: deltat = 0.001
# the emission rate, 20Hz
In [65]: alpha = 20
# 1000 random numbers
In [66]: rvs = uniform.rvs(1000)
```

```
# a look at the 1st 20 random variates
In [67]: rvs[:20]
Out[67]:
array([ 0.71167172, 0.01723161, 0.25849255, 0.00599207, 0.58656146,
        0.12765225, 0.17898621, 0.77724693, 0.18042977, 0.91935639,
        0.97659579, 0.59045477, 0.94730366, 0.00764026, 0.12153159,
        0.82286929, 0.18990484, 0.34608396, 0.63931108, 0.571991751)
# we simulate an emission when the random number is less than
\# p(Delta t) = alpha * deltat
In [84]: emit = rvs < (alpha * deltat)</pre>
# there were 3 emissions in the first 20 observations
In [85]: emit[:20]
Out[85]:
array([False, True, False, True, False, False, False, False, False,
       False, False, False, False, False, False, False, False,
       False, False], dtype=bool)
```

The waiting times between the emissions should follow an exponential distribution (see scipy.stats.expon) with a mean of $1/\alpha$. In the exercise below, you will generate a long array of emissions, compute the waiting times between emissions, between 2 emissions, and between 10 emissions. These should approach an 1st order gamma (aka exponential) distribution, 2nd order gamma, and 10th order gamma (see scipy.stats.gamma). Use the probability density functions for these distributions in scipy.stats to compare your simulated distributions and moments with the analytic versions provided by scipy.stats. With 10 waiting times, we should be approaching a normal distribution since we are summing 10 waiting times and under the central limit theorem the sum of independent samples from a finite variance process approaches the normal distribution (see scipy.stats.norm). In the final part of the exercise below, you will be asked to approximate the 10th order gamma distribution with a normal distribution. The results should look something like those in Figure 2.

Listing 7.2

```
. . . .
Illustrate the connections bettwen the uniform, exponential, gamma and
normal distributions by simulating waiting times from a radioactive
source using the random number generator. Verify the numerical
results by plotting the analytical density functions from scipy.stats
import numpy
import scipy.stats
from pylab import figure, show, close
# N samples from a uniform distribution on the unit interval. Create
# a uniform distribution from scipy.stats.uniform and use the "rvs"
# method to generate N uniform random variates
N = 100000
uniform = XXX # the frozen uniform distribution
uninse = XXX # the random variates
# in each time interval, the probability of an emission
rate = 20. # the emission rate in Hz
dx = 0.001 # the sampling interval in seconds
```

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```
t = numpy.arange(N) *dx # the time vector
# the probability of an emission is proportionate to the rate and the interval
emit\_times = XXX
# the difference in the emission times is the wait time
wait\_times = XXX
# plot the distribution of waiting times and the expected exponential
# density function lambda exp( lambda wt) where lambda is the rate
# constant and wt is the wait time; compare the result of the analytic
# function with that provided by scipy.stats.exponential.pdf; note
# that the scipy.stats.expon "scale" parameter is inverse rate
# 1/lambda. Plot all three on the same graph and make a legend.
# Decorate your graphs with an xlabel, ylabel and title
fig = figure()
ax = fig.add_subplot(111)
p, bins, patches = XXX # use ax.hist
11, = ax.plot(bins, XXX, lw=2, color='red') # the analytic result
12, = ax.plot(bins, XXX,
                                             # use scipy.stats.expon.pdf
        lw=2, ls='--', color='green')
ax.set_xlabel('waiting time')
ax.set_ylabel('PDF')
ax.set_title('waiting time density of a %dHz Poisson emitter'%rate)
ax.legend(XXX, XXX) # create the proper legend
# plot the distribution of waiting times for two events; the
# distribution of waiting times for N events should equal a N-th order
# gamma distribution (the exponential distribution is a 1st order
# gamma distribution. Use scipy.stats.gamma to compare the fits.
# Hint: you can stride your emission times array to get every 2nd
# emission
XXX
# plot the distribution of waiting times for 10 events; again the
# distribution will be a 10th order gamma distribution so plot that
# along with the empirical density. The central limit thm says that
# as we add N independent samples from a distribution, the resultant
# distribution should approach the normal distribution. The mean of
\sharp the normal should be N times the mean of the underlying and the
# variance of the normal should be 10 times the variance of the
# underlying. HINT: Use scipy.stats.expon.stats to get the mean and
# variance of the underlying distribution. Use scipy.stats.norm to
# get the normal distribution. Note that the scale parameter of the
# normal is the standard deviation which is the square root of the
# variance
XXX
```

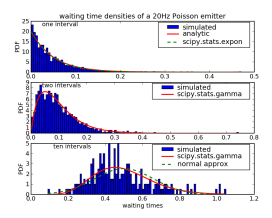


FIGURE 2.